

Title Page

Manuscript title The first description and quantitative assessment of the
conjunctival microcirculatory profile using a smartphone

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26 **Abstract**

27 **Purpose** The conjunctival microcirculation is a readily-accessible vascular bed
28 for quantitative haemodynamic assessment and has been studied previously using a
29 digital charge-coupled device (CCD). Smartphone video imaging of the conjunctiva,
30 and haemodynamic parameter quantification, represents a novel approach. We
31 report the feasibility of smartphone video acquisition and subsequent haemodynamic
32 measure quantification via semi-automated means.

33 **Methods** Using an Apple iPhone 6s and a Topcon SL-D4 slit-lamp
34 biomicroscope, we obtained videos of the conjunctival microcirculation in 4 fields of
35 view per patient, for 17 low cardiovascular risk patients. After image registration and
36 processing, we quantified the diameter, mean axial velocity, mean blood volume
37 flow, and wall shear rate for each vessel studied. Vessels were grouped into
38 quartiles based on their diameter i.e. group 1 ($<11\mu\text{m}$), 2 ($11\sim16\mu\text{m}$), 3 ($16\sim22\mu\text{m}$)
39 and 4 ($>22\mu\text{m}$).

40 **Results** From the 17 healthy controls (mean QRISK3 6.6%), we obtained
41 quantifiable haemodynamics from 623 vessel segments. The mean diameter of
42 microvessels, across all sites, was $18.23\mu\text{m}$ (range $6.6\sim39.2\mu\text{m}$). Mean axial velocity
43 was 0.49mm/s (range $0.12\sim0.79\text{mm/s}$) and there was a modestly positive correlation
44 ($r\ 0.404$) seen with increasing diameter, best appreciated when comparing group 4
45 to the remaining groups ($p<0.0001$). Blood volume flow (mean 109.718pl/s , range
46 $11.28\sim502.19\text{pl/s}$) was strongly correlated with increasing diameter ($r\ 0.967$,
47 $p<0.0001$) and wall shear rate (mean 182.81s^{-1} , range $55.11\sim546.69\text{s}^{-1}$) negatively
48 correlated with increasing diameter ($r\ -0.823$, $p<0.0001$).

Conclusions We, for the first time, report the successful assessment and quantification of the conjunctival microcirculatory haemodynamics using a smartphone-based system.

Manuscript

I. Introduction

Cardiovascular disease (CVD) is a leading cause, globally, of mortality and morbidity while also being associated with a significant economic burden on health services¹.

CVD is caused by physiological changes and endothelial dysfunction, resulting in atherosclerosis, and it is accepted that these changes manifest earliest in the microcirculatory networks within the body². Microcirculatory disease typically

commences with endothelial dysfunction which may be clinically silent and, thus, precede the onset of symptoms³ or the occurrence of a major adverse

cardiovascular event (MACE) e.g. myocardial infarction (MI) or cerebrovascular accident (CVA). Microvascular dysfunction is associated with increased mortality⁴

and thus the study of microcirculations may provide a potential tool in disease

screening, staging and management. Imaging of systemic microcirculations has

been applied to and, in certain disease subsets, is used in every day current practice in assessing disease progression e.g. the retinal microcirculation in the assessment

of diabetes mellitus, systemic hypertension, and sickle cell disease^{5,6,7,8}. The

sublingual mucosa and the skin also represent accessible sites in which the

microcirculation has been studied by videomicroscopy⁹.

The anterior segment of the eye contains the conjunctival microvasculature, a

readily-accessible heterogeneous network of arterioles and venules adjacent to the

limbal microcirculation, which gains its supply from the anterior ciliary branch of the

74 ophthalmic artery¹⁰. The conjunctival microvasculature allows for both non-invasive
75 assessment of erythrocyte movement, and quantification of key vascular
76 physiological parameters e.g. vessel width, blood flow axial velocity and blood flow
77 rate¹¹.

78 The objective of this study was to evaluate the feasibility of assessing the
79 conjunctival microcirculation using our novel combination of a smartphone and slit-
80 lamp biomicroscope. We aimed to develop an operator-friendly, pragmatic, safe and
81 effective means of assessing this heterogeneous circulation, in addition to the
82 quantification of the haemodynamic physiological parameters seen within a
83 microcirculation.

84 A few groups have reported semi-automated or automated image analysis
85 algorithms to assess the conjunctival microcirculation, using a slit lamp
86 biomicroscope and a digital charge-coupled device (CCD) camera for image
87 acquisition^{12,13,14,15,16,17}. Using such systems, the conjunctival microcirculation has
88 been studied in patients with hypertension, diabetic retinopathy, and patients after
89 ischaemic stroke^{18, 19, 20}. In addition, one group has reported the application of such
90 methods in patients of varying predictive cardiovascular risk, assessed by the
91 Framingham risk score²¹.

92 Smartphone technology allows for remote monitoring and screening of many
93 prevalent cardiovascular conditions, for example atrial fibrillation, and represents an
94 important component of future healthcare and cardiovascular practice²². The
95 literature is scarce regarding smartphone use to assess microcirculatory
96 haemodynamics but the application of smartphone photography of the fundus has
97 been reported in diabetic and hypertensive patients^{23, 24, 25}. [There are some studies](#)
98 [describing smartphone-led image analysis of the conjunctiva in the assessment of](#)

99 patients with anaemia^{26,27} and, also, quantification of conjunctival “redness” i.e.
100 hyperaemia²⁸. In addition, the smartphone-based biometric has been studied on the
101 visible vascular patterns on whites of the eye²⁹ but, at this time, there are no studies
102 that describe the assessment or quantification of conjunctival haemodynamics using
103 a smartphone and slit-lamp combination. ↓

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105 **II. Materials and Methods**

106 **A. Subjects**

107 This research study was approved by the Research and Development review boards
108 of the Ulster University (UU) and the Belfast Health and Social Care Trust (BHSCT).
109 All subjects were provided with verbal and written information, prior to study
110 enrolment, in accordance with the Declaration of Helsinki. Exclusion criteria included
111 inability to consent, prior myocardial infarction (MI), uncontrolled systemic
112 hypertension, recent history of conjunctival inflammation, prior refractive surgery,
113 used ocular medications (other than artificial tears) and current use of contact
114 lenses.

115 We recruited 17 healthy volunteers to this feasibility study. The mean age for the
116 population studied was 52.5 ±10.3years, IQR 15 years. Sex distribution was roughly
117 equal with 9 (53%) males and 8 females (47%). No patients had a history of prior MI,
118 cerebrovascular accident (CVA), or diabetes mellitus. The well-validated QRISK 3
119 (<https://qrisk.org/three/>) score algorithm was used to estimate each volunteer's 10-
120 year risk of future heart attack or stroke. The QRISK 3 algorithm is based on the
121 presence/lack of specific risk factors for CVD e.g. smoking, diabetes mellitus,
122 hypertension, family history angina, chronic kidney disease, age, sex, body mass
123 index, history of atrial fibrillation, use of regular steroid tablets, presence of chronic

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inflammatory disease, and cholesterol profile. The mean QRISK 3 score was 6.6
 ±9%, IQR 6.9%, which correlates with a “low-risk” population (<10%)³⁰. Table 1 is a
 summary of the baseline demographics and clinical observations for the study group.

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	Number n=17
Male sex, n (%)	9 (53.0)
Age, years ±SD	52.5 ±10.3
QRISK 3 score, % ±SD	6.6 ±9
Systolic blood pressure, mmHg ±SD	125 ±22
Diastolic blood pressure, mmHg ±SD	77 ±12
Heart rate, bpm ±SD	70 ±9
Prior MI/CVA/Diabetes mellitus	0

Table 1. Baseline characteristics of the study group (n=17) with continuous variables expressed using their mean and standard deviation. Categorical variables have been expressed as a number and percentage of the total within that variable.

B. Image Acquisition

Image acquisition was achieved via two main hardware components. Firstly, primary illumination and magnification of the ocular vascular structure was achieved using a conventional slit lamp biomicroscope, Topcon SL-D4 (Topcon Medical Systems Inc., USA), capable of providing a maximum magnification of 40x. Secondly, images provided by the slit lamp biomicroscope were further magnified and stored using a smartphone camera. The smartphone used in the system is an Apple iPhone 6s (Apple, Inc., USA). A number of video record settings were tested and the optimal configuration set at a resolution of 1920 x 1080 pixels, captured at 60 frames per second. The iPhone video recorder is capable of providing a further magnification of 3x. Coupling of the smartphone to the eyepiece of the slit lamp biomicroscope was

148 achieved using a bespoke adapter developed by Zarf Enterprises (Zarf Enterprises.,
149 USA). Smartphone cameras typically give very little control over camera properties
150 (focus, ISO, shutter speed, aperture) due to an emphasis on ease-of-use for
151 everyday consumers, while also generating compressed video files (h.264
152 compression in the case of the iPhone 6s). To help overcome these issues we
153 captured our data using a third-party application "ProMovie Recorder"
154 (www.promovieapp.com). We used constant settings for all images (iso/shutter
155 speed/ focus/ exposure) and used the maximum compression bit-rate available to
156 reduce compression artefacts. The video zoom setting was locked at 2x, providing a
157 1:1-pixel mapping of the camera sensor at 1080p resolution and thus avoiding
158 interpolation artefacts. To obtain an accurate pixel to mm conversion factor we
159 calibrated the system using a digital caliper and 1mm microscope calibration reticle,
160 deriving a conversion factor of 552 ± 22.6 pixels/mm. We obtained one video (5-15s)
161 from 4 distinct field of views i.e. medial and temporal conjunctiva in both eyes. Fig.1.
162 To reduce eye motion and blinking we used an external fixation target as a focal
163 point for each patient. We acquired only 4 videos (5-15s) per patient to minimise the
164 risk of potential adverse effects, e.g. slit-lamp light exposure. There were no reported
165 adverse effects at the time of, or after, image acquisition. Patients were imaged in
166 the same clinical room under constant temperature and lighting settings.



Fig. 1. Two fields of view (FOV) for the left eye of a healthy subject, with the medial and lateral FOV being labelled (red arrows) the left nasal (LN) and left temporal (LT) respectively.

C. Image Processing

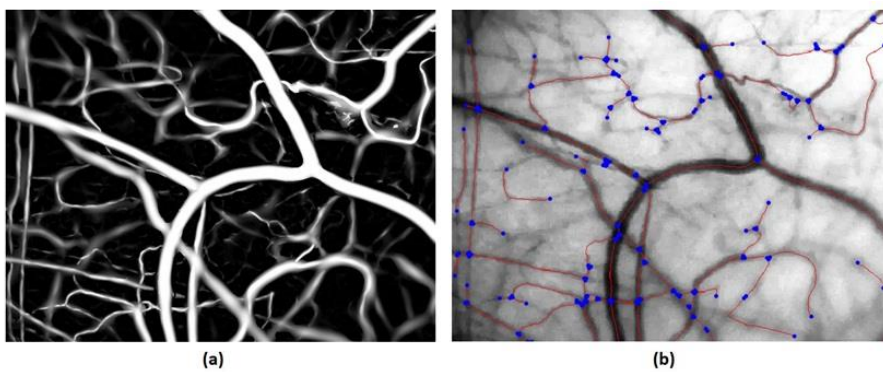
1. Pre-Processing and Vessel Segmentation

An initial pre-processing procedure was carried out for each video file. Firstly, the longest stable sequence of frames was manually selected on the basis of the vasculature being in focus, there being no blinking or large sudden movements of the eye, and the FOV not drifting by more than ~25% of the width of the frame. Next the green channel, which gave the highest vessel contrast, was extracted and information from the red channel used to correct for uneven illumination through subtraction. The sharpest frame in the sequence was then selected as a reference frame and all other frames registered to it through an affine registration procedure³¹ with a single composite image generated by averaging all registered frames. After applying a “vessel enhancement filter”³² (Fig.2 (a)), a binary map of the conjunctival

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187 vasculature and corresponding centrelines were obtained via standard skeletisation
 188 techniques. Finally, the connected vessel network was broken into individual vessel
 189 segments (Fig.2 (b)) by setting the branch points' neighbouring pixels to zero, and
 190 centreline segments, containing more than 30 pixels, selected for further
 191 assessment.



192 (a) (b)
 193 Fig. 2. Microvascular network after image processing: (a) the vessel network after
 194 filtering; (b) the vessel centreline (in red) and intersection points (in blue) overlaid on
 195 the mean of vessel images.

196 2. Vessel Diameter (D)

197 The Euclidean Distance Transform (EDT) was proposed for vessel diameter
 198 estimation, which is easier to implement in comparison to the commonly used
 199 method via full width at half maximum (FWHM). The value at each pixel of EDT was
 200 calculated based on the Euclidean distance between the pixel and its nearest
 201 nonzero pixel in the binary vessel image. The centreline of the vessel was used to
 202 obtain the central EDT values and thus the radius along the vessel axis. The
 203 average of diameters along the vessel length provided the final vessel width
 204 estimation. An example based on simulation is illustrated in Fig.3.

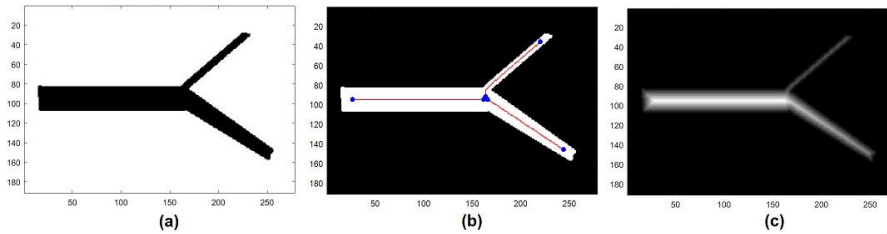


Fig. 3. Simulation for vessel diameter estimation: (a) three vessels are generated with mean diameter 25.3 pixels, 16.5 pixels, and 8.3 pixels, respectively; (b) the vessel centreline, end points and branch points overlaid on the binary vessel image; (c) EDT of the binary vessel image. The mean of estimated diameters via EDT are 25.9 pixels, 16.6 pixels, and 8.6 pixels, respectively.

Given the complex and heterogeneous distribution of conjunctival microvessels, we applied a grouping classification to our results, described in previous work, based on vessel D i.e. group 1 ($<11\mu\text{m}$), group 2 ($11\text{-}16\mu\text{m}$), group 3 ($16\text{-}22\mu\text{m}$) and group 4 ($>22\mu\text{m}$)¹¹.

3. Axial velocity (V_a)

The blood flow V_a in a single vessel segment was estimated based on the spatial-temporal image (STI), with the change in intensity in STI reflecting erythrocyte movement through the vessel. Since STI signal is the one dimension of space plus time, i.e., a 1D+T signal, a novel approach based on spatial temporal 1D+T continuous wavelet transform (1DTCWT) is proposed for V_a estimation. The CWT method has been used previously as a spatiotemporal filter for motion capture of 1D+T signals for moving target tracking and parameter calculation³³, but not yet exploited in microvascular blood flow velocity estimation. Firstly, 2D fast Fourier transform (FFT) is performed for STI. The velocity vector space is defined and

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1DTCWT is then run at each time interval. The energy is subsequently calculated based on the 1DTCWT output. The velocity is obtained by searching the maximum energy point as shown in Fig.4. The average of the absolute velocity across all frames was used as the final estimation of V_a . The method was programmed in MATLAB2017 together with an open source implementation of CWT³⁴

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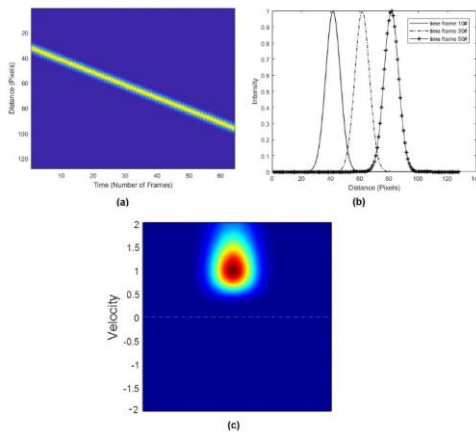


Fig.4. Simulation for velocity estimation based on 1DTCWT. (a) synthetic STI generated by shifting Gaussian signal with speed of 1 pixel/frame; (2) plot of signals at the 10th, 30th and 50th frames, which shows the Gaussian signal shifting in distance; (c) a colour spectrum map via 1DTCWT shows the velocity is corresponding to the maximum of the energy (at 1 pixel/frame).

4. Blood flow (Q) and wall shear rate (WSR)

Using the measurements for D and V_a , we calculated Q and WSR using previously described methods^{11,12}. Q provides key information regarding the architecture and function of the vascular system, whereas WSR is the blood velocity at a specific wall

242 position, within a vessel, and represents a surrogate for the pressure exerted by
243 blood within its' respective transport vessel ^{35, 36, 37}

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245 5. Statistical analysis

246 For statistical analysis SPSS for Apple iOS version 25 (property of IBM) and R
247 version 3.5.3 (www.r-project.org) were used. Continuous variables were described
248 using the mean, standard deviation of the mean and interquartile range (IQR) for the
249 variable. Categorical variables were described as a number and percentage of the
250 total category number to which the variable belonged. Sample origin, distribution and
251 variance were assessed by non-parametric ANOVA (Kruskal-Wallis test). Correlation
252 analysis (Spearman rank), with a Loess regression fit, was applied to assess
253 relationships between D and independent variables, principally Va, Q and WSR.
254 Non-parametric ANOVA (Kruskal-Wallis) with or without Dunn's post-hoc tests was
255 used to compare D, Va, W, and WSR by vessel width group, with the tests being
256 conducted separately across site, i.e., left/right nasal and temporal, or for all sites
257 merged.

258 **III. Results**

259 For the 17 healthy patients studied, using our semi-automated approach, we were
260 able to obtain repeated measurements in 623 vessel segments (mean 37 segments
261 per patient), hereafter referred to as "microvessels", which exhibited observable flow.
262 The mean diameter (D) of microvessels, across all sites, was 18.2 μ m (range 6.6-
263 39.2 μ m). Group 4 (>22 μ m) microvessels were measured most frequently, with group
264 1 (<11 μ m) being the least commonly encountered i.e. 295 vs 64 microvessels
265 respectively. Mean Va was 0.49mm/s (range 0.12-0.79mm/s), Q 109.72pl/s (range

269 11.28-502.19pl/s) and WSR ranged between 55.11-546.69s⁻¹, with a mean WSR of
 270 182.81s⁻¹. The mean and SD of all microvessel conjunctival haemodynamic
 271 parameters are illustrated in Table 2. Statistical comparisons for Va, Q and WSR
 272 were made within the vessel groups. There was a statistically significant increase in
 273 Q for increasing diameter size (p<0.0001), with a statistically significant inverse
 274 correlation between WSR and increasing diameter size (p<0.0001). Va tended to
 275 increase with increasing microvessel diameter and was significantly elevated in
 276 group 4 (>22µm) vessels, compared to the remaining three groups (p<0.0001).
 277
 278

Group D µm	No. vessels N=623	D (µm)	Va (mm/s)	Q (pl/s)	WSR (s ⁻¹)
<11 Group 1	64	9.1 ±2.8	0.45 ±0.05	23.65 ±2.96	332.75 ±60.75
11~16 Group 2	113	13.44±3.7	0.44 ±0.06	46.81 ±8.02	200.19 ±32.89
16~22 Group 3	151	19.2 ±3.5	0.47 ±0.06	97.13 ±17.21	136.67 ±20.35
>22 Group 4	295	26.9 ±2.7	0.56 ±0.09	224.45 ±66.35	115.27 ±17.7
			p<0.0001	p<0.0001	p<0.0001
	Mean	18.2	0.485	109.718	182.81
	Range	6.6-39.2	0.12-0.79	11.28-502.19	55.11-546.69
	Interquartile range (IQR)	12.74-24	0.42-0.55	39.86-161	116.33- 221.72

Table 2. Summary of haemodynamic measures D, Va, Q and WSR based on the vessel diameter groups (1-4).

Across site (field of view) comparisons were made with the haemodynamic measures. Q and WSR did not statistically differ between the 4 image fields. There was a statistically higher Va noted in the right nasal (RN) hemisphere compared to the left nasal (LN, ($p = 0.0003$)), for which the clinical significance is unknown and may require further exploration. The relationship between the haemodynamic measures and similarities for each field of view is shown in Fig.5. Note the elevated Va in the RN FOV, compared to the other FOVs, as before.

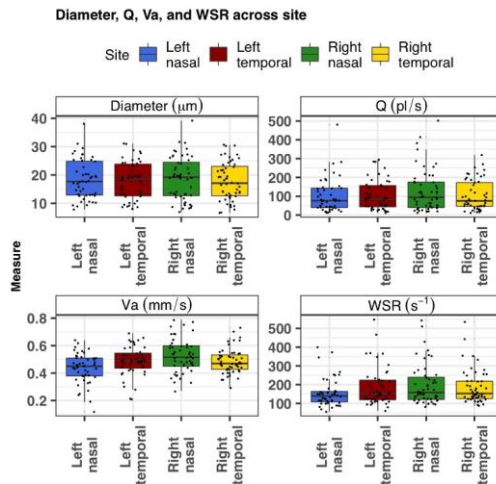
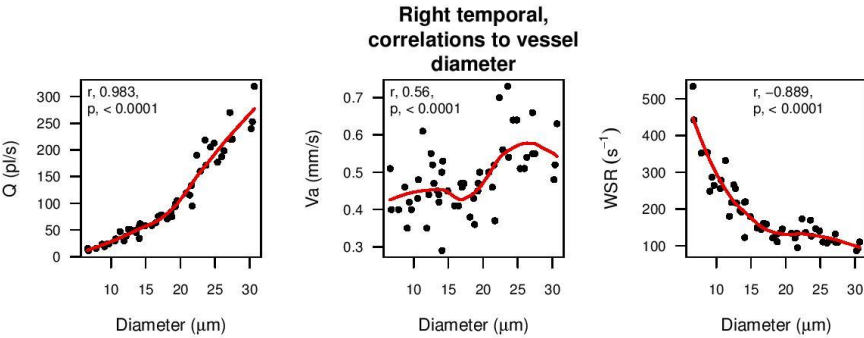


Fig.5. Summary of diameter D (μm), Va (mm/s), Q (pl/s) and WSR (s^{-1}), for each field of view i.e. left nasal (LN), left temporal (LT), right nasal (RN) and right temporal (RT).

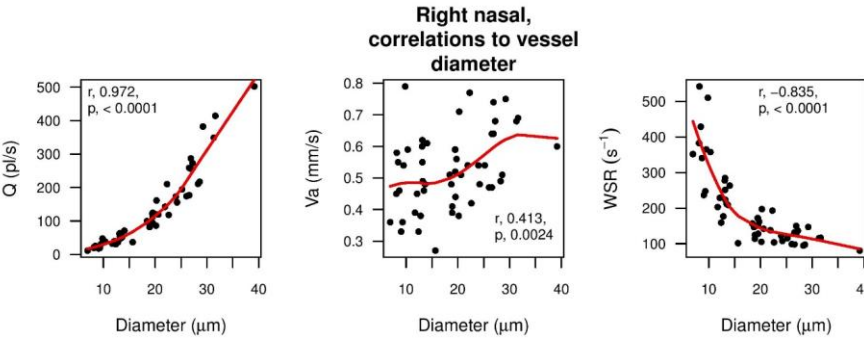
The correlation, expressed via the correlation coefficient (r) and the best fit trend line, between increasing microvessel diameter and the haemodynamic measures Va, Q and WSR were consistent across the 4 fields of view, which are individually

296 illustrated in Fig.6a-d.

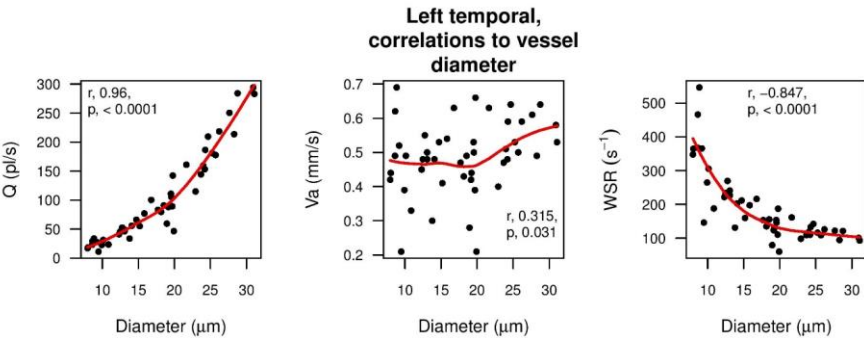
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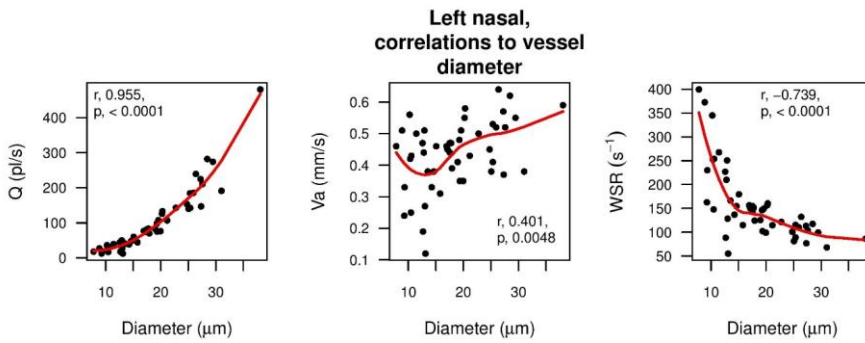


Fig.6. Correlation plots between microvessel diameter D (μm) vs V_a (mm/s), Q (pl/s) and WSR (s^{-1}) for each field of view ((a) RT, (b) RN, (c) LT, (d) LN).

A summary of the correlations between microvessel D and the quantified haemodynamic measures are illustrated in Fig.7. demonstrating the strong overall linear correlation with Q and WSR (r 0.967, r -0.823 respectively). A modest correlation was seen for V_a (r 0.404).

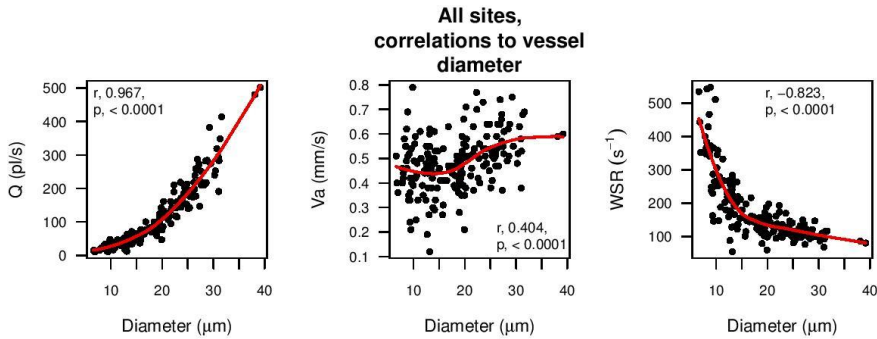


Fig.7. Correlation plots between microvessel diameter D (μm) vs V_a (mm/s), Q (pl/s) and WSR (s^{-1}) across all sites.

The correlations between increasing vessel diameter and V_a , Q , and WSR are in keeping with that reported in previous work^{11,12}, whereby similar fluid dynamics and microvascular relationships have been observed.

IV. Discussion

The conjunctival microcirculation represents a readily-accessible vascular network for non-invasive assessment. Physiological measures in the conjunctival microcirculation display the same trends and correlations as they do elsewhere in the circulation and, based on this rationale, may represent a key microcirculation that could be assessed in the evaluation of circulatory health and, if so, correlated with risk. Correlations between cardiovascular risk estimation and quantitative conjunctival haemodynamic measures, namely velocity and blood flow, were demonstrated in previous work²¹.

In recent years, there have been several reports regarding the clinical utility of conjunctival microcirculatory study. Conjunctival haemodynamic assessment has extended to patients with diabetes mellitus, in correlation with diabetic retinopathy status, with differences between V_a , Q and WSR being observed for differing grades of retinopathy¹⁹. Quantitative assessment of the conjunctival haemodynamics was, also, evaluated in patients with ischaemic unilateral stroke and V_a was found to be significantly lower in the ipsilateral eye to the stroke compared to the contralateral eye, demonstrating the physiological relationship shared by the internal carotid arterial system and the conjunctival microcirculation²⁰.

We have described the application of smartphone technology, combined with a slit-lamp, in the quantitative assessment of conjunctival haemodynamics, namely D , V_a , Q and WSR. With our approach, we have demonstrated the feasibility of obtaining haemodynamic results, similar to the correlations and trends described elsewhere by other groups using a digital charged coupled camera. We have done so, though, using a smartphone which served as an efficient, pragmatic and reliable means of

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339 acquiring the conjunctival images for subsequent analysis. Our system performed as
340 well as the more complex and time-consuming CCD devices and represents a
341 potential major advancement within the scope of conjunctival microcirculation
342 assessment. Our biomicroscope/smartphone apparatus and post-capture analysis is
343 validated by comparison to results obtained previously. We obtained a mean
344 diameter of 18.2µm (range 6.6-39.2µm) in 623 microvessels, selected manually
345 according to the quality of STI, on post-processed images and these results are
346 similar to, and within range, of that reported by other groups¹¹. The strong
347 positive/negative correlation between microvessel diameter (D) and blood flow (Q)/
348 wall shear rate (WSR), reported in the present work, is in keeping with that found in
349 other studies^{11, 12, 13}. We did not find as strong a correlation for axial velocity (Va) and
350 diameter (D) ($r = 0.404$), compared to that observed for blood flow (Q) and wall shear
351 rate (WSR). Statistical significance, though, was observed for group-4 vessels and
352 their associated Va, compared to groups 1-3 ($p < 0.0001$).

353 Combined smartphone and slit-lamp based quantitative assessment has been
354 demonstrated in this present work and it is feasible that it could be of potential future
355 application in the assessment of cardiovascular health. We studied a “low-
356 cardiovascular risk” patient group, as evidenced by a mean QRISK 3 score of 6.6%.
357 QRISK 3 is a well-validated 10-year cardiovascular risk assessment, with the largest
358 sample size of contemporary cardiovascular estimation systems, implemented within
359 major European guidelines³⁰.

360 We acknowledge certain limitations of our study. We, similar to other feasibility
361 studies¹³, have reported results for all visible microvessels without separating
362 arterioles and venules. The feasibility of artery-vein classification, using our
363 approach, in the conjunctiva requires further exploration, which we intend to pursue.

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370 In addition, cardiac-gated haemodynamic measures, primarily end-systolic and end-
371 diastolic measures using conjunctival vessel pulse waveform characteristics, have
372 been reported previously and could be of potential use in future clinical application
373 with certain cardiovascular disease subsets³⁵. A key aim of our future work is to
374 implement and validate a fully automated smartphone-based approach to remove
375 potential human error, promote consistency, and improve the efficiency of the
376 examination. By quantifying the conjunctival haemodynamics our method potentially
377 allows the inexpensive assessment of patients with established cardiovascular and
378 systemic disease, with promise for improving the diagnosis, risk stratification and,
379 potentially, evaluating disease status and treatment modification of cardiovascular
380 disease(s). The addition of smartphone technology, with it's APP versatility, wealth of
381 data management, and computerised machine learning algorithms, to a slit-lamp
382 biomicroscope potentially modernises the assessment of the conjunctival
383 microcirculation.

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384 V. Conclusion

385
386 We have described, for the first time, the successful measurement of dynamic
387 microcirculatory haemodynamic measures using smartphone technology combined
388 with a slit-lamp biomicroscope. Our semi-automated method found a positive linear
389 relationship between increasing microvessel diameter (D) and blood flow (Q). An
390 inverse relationship was observed for wall shear rate WSR, a direct surrogate of
391 WSS. These findings corroborate prior ones, for the same haemodynamic measures,
392 reported by groups using a CCD camera for image acquisition, and support the
393 feasibility of our smartphone-derived approach. Image acquisition was performed
394 without clinical complication in a group of patients with low cardiovascular risk. The

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401 ease and speed with which images were reliably acquired holds promise for the
402 future clinical application of this smartphone-based conjunctival microcirculatory
403 assessment model.

404

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406 **VI. Acknowledgements**

407 This project was funded by Northern Ireland Chest Heart and Stroke (NICHHS), the
408 Ulster University and the Heart Trust fund, Royal Victoria Hospital, Belfast, United
409 Kingdom.

410 **VII. Disclosure/Conflict of interests**

411 The authors, collectively, have no conflicts of interest or anything to disclose with
412 respect to this original research manuscript.

413

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